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                               SUBSTITUTE SHEET (RULE 26)
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<213> Homo sapiens - human MDC

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Glu Tyr Phe Tyr Thr Ser Gly Lys Cys Ser Asn Pro Ala Val Val Phe
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Val Arg Glu Tyr Ile Asn Ser Leu Glu/Met Ser
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Gln Thr Lys Arg Ser Lys Gln Val Cys Ala Asp Pro Ser Glu Ser Trp
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Cys Cys Phe Ser Tyr Thr Ser Arg Gln Ile Pro Gln Asn Phe Ile Ala
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                                     20
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Cys Tyr Arg Asn Thr Ser Ser Me Cys Ser Asn Glu Gly Leu Ile Phe

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<223> The amino acid at position 30 is independently selected from the group consisting of tyr, ser, lys, arg, his, aspartate, glutamate, asn, gln, and cys

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<223> The amino acid at position 50 is independently selected from the group consisting of glu, lys, arg, his, gly, and ala

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<223> The amino acid at position \$9 is independently selected from the group consisting of trp, ser, lys, arg, his, aspartate, glutamate, asn, gln, and cys

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<223> The amino acid at position 60 is independently selected from the group consisting of val. ser, lys, arg, his, aspartate, glutamate, asn, gln, and cys

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aga aa i tac Tyr	aaag Met 1 agc Ser	caa g aac g Asn i aat Asn	getg ecc : Pro ' tac Tyr	acg grant tat	gat Asp : Ctg Leu	ata gata la	geoca gra gaa Glu	gat a Asp agt ser	acc a Thr 1 atc Ile 25	acc (Thr 10 10 ccc	gagg etc g Leu i aag Lys	gat	gaa ; Slu ; tgc Cys	agc a Ser I acc Thr	ttaa ita ile 15 aaa Lys	227
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1	

Tyr Tyr Ala Ala Asp Gln Trp Val Phe Gly Leu Gly Leu Cys Lys Met 100 105 110

Ile Ser Trp Met Tyr Leu Val Gly Phe Tyr Ser Gly Ile Phe Phe Val

Met Leu Met Ser Ile Asp Arg Tyr Leu Ala Ile Val His Ala Val Phe 130 135 ___ 140

Ser Leu Arg Ala Arg Thr Leu Thr Tyr Gly Val Ile Thr Ser Leu Ala 145 150 155 160

Thr Trp Ser Val Ala Val Phe Ala Ser Leu Pro Gly Phe Leu Phe Ser

Thr Cys Tyr Thr Glu Arg Asn His Thr Tyr Cys Lys Thr Lys Tyr Ser 180 185 190

Leu Asn Ser Thr Thr Trp Lys Val Leu Ser Ser/Leu Glu Ile Asn Ile
195 200 205

Leu Gly Leu Val Ile Pro Leu Gly Ile Met Leu Phe Cys Tyr Ser Met 210 215 220

Ile Ile Arg Thr Leu Gln His Cys Lys Asn Glu Lys Lys Asn Lys Ala
225 230 235 240

Val Lys Met Ile Phe Ala Val Val Leu Phe Leu Gly Phe Trp Thr
245 250 255

Pro Tyr Asn Ile Val Leu Phe Leu/Glu Thr Leu Val Glu Leu Glu Val
260 / 265 270

Leu Gln Asp Cys Thr Phe Glu Arg Tyr Leu Asp Tyr Ala Ile Gln Ala 275 /280 285

Thr Glu Thr Leu Ala Phe Vai His Cys Cys Leu Asn Pro Ile Ile Tyr 290 / 295 300

Phe Phe Leu Gly Glu Lys Phe Arg Lys Tyr Ile Leu Gln Leu Phe Lys 305 310 315 320

Thr Cys Arg Gly Leu Phe Val Leu Cys Gln Tyr Cys Gly Leu Leu Gln 325 330 335

Ile Tyr Ser Ala Asp Thr Pro Ser Ser Ser Tyr Thr Gln Ser Thr Met
340 345 350

Asp His Asp Leu His Asp Ala Leu 355 / 360

<210> 35 <211> 1784

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•		,	7	
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Met Se	er Asn Leu Arg Val -20

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Met Ser Asn Leu Arg Val Pro Leu Leu Val Ala Leu Val Leu Leu Ala
-20 -15 -10

gtg gca att cag acc tct gat gca ggt ccc tat gg/t gcc aat gtg gaa 96
Val Ala Ile Gln Thr Ser Asp Ala Gly Pro Tyr Gly Ala Asn Val Glu
-5 -1 1 5

gac agt atc tgc tgc cag gac tac atc cgt cac cct ctg cca tca cgt

Asp Ser Ile Cys Cys Gln Asp Tyr Ile Arg H.s Pro Leu Pro Ser Arg

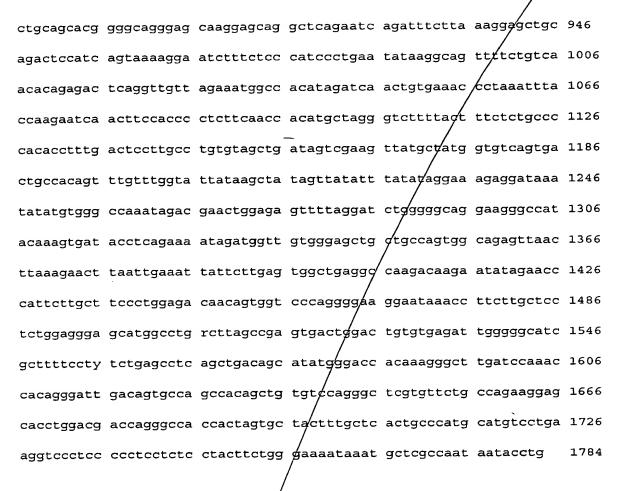
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tta gtg aag gag ttc ttc tgg acc tca aaa tcc tgc cgc aag cct ggc
Leu Val Lys Glu Phe Phe Trp Thr Ser Lys Ser Cys Arg Lys Pro Gly
25 30 35 40

gtt gtt ttg ata acc gtc aag aac cga gat atc tgt gcc gat ccc agg 240
Val Val Leu Ile Thr Val Lys Asn Arg Asp Ile Cys Ala Asp Pro Arg
45 50 55

cag gtc tgg gtg aag aag cta ctc/cat aaa ctg tcc tagggaggag 286
Gln Val Trp Val Lys Lys Leu Leu His Lys Leu Ser
60 65

gacctgatga ccatgggtct ggtgggtcc agggaggctc agcaagccct attetetge 346 catteragea agageettge caacgacgec acetttacte acetecatee eetgggetgt 406 cactetgtca ggetetggte eetetacete eetetatee ettecagett atececette 466 aatgtggcag etgggaaaca catteaggee ageettacee aatgeetaet eeecaatget 526 ttagatgaga ecagegteet tgttttgatg eeetgateet atgatgeett eeecaateee 586 ageettggee eeettetett ettgeatgta gggaaggeee ataggttea aatatgtget 646 aeetaettee ettetetggg ggttetaata eeeageatgt tttteetget geaggeacet 706 atecagtgee acacacetee eaagttteta teagteecag tgggcateea ecaageecea 766 aaetteagae tteettggee teeacetaet eteagtagaa ttetgggagt tteaggetgg 826 teeaceagge eeeeeagggt taggeeaagg teeceaceag ageteeteet gttettggt 886



<210> 36 <211> 92 <212> PRT

<213> murine MDC

<400> 36

Met Ser Asn Leu Afg Val Pro Leu Leu Val Ala Leu Val Leu Leu Ala

Val Ala Ile Gln Thr Ser Asp Ala Gly Pro Tyr Gly Ala Asn Val Glu
-5 -1 1 5

Asp Ser Ile Cys Cys Gin Asp Tyr Ile Arg His Pro Leu Pro Ser Arg
10 20

Leu Val Lys Glu Phe Phe Trp Thr Ser Lys Ser Cys Arg Lys Pro Gly 25 30 35 40

Val Val Leu Ile Thr Val Lys Asn Arg Asp Ile Cys Ala Asp Pro Arg
45 50 55

Gln Val Trp Va Lys Lys Leu Leu His Lys Leu Ser

<210> 37 <211> 958	
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<220> <221> CDS <222> (1)(243)	
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ggt gcc aat gtg gaa gac agt atc tgc tgc cag gac tac atc cgt cac 96 Gly Ala Asn Val Glu Asp Ser Ile Cys Cys Gln Asp Tyr Ile Arg His 5 10 15	6
cct ctg cca cca cgt ttc gtg aag gag ttc tac tgg acc tca aag tcc Pro Leu Pro Pro Arg Phe Val Lys Glu Phe Tyr Trp Thr Ser Lys Ser 20 25 30 35	44
tgc cgc aag cct ggc gtc gtt ttg ata acc atc aag aac cga gat atc Cys Arg Lys Pro Gly Val Val Leu Ile Thr Ile Lys Asn Arg Asp Ile 40 45 50	92
tgt gct gac ccc ang atg ctc tgg gtg aag aag ata ctc cac aag ttg Cys Ala Asp Pro Xaa Met Leu Tap Val Lys Lys Ile Leu His Lys Leu 55 60 65	40
gcc tagggagaag ggcctgatga ccadgggtct ggtgtctcca caaggctcag 29	93
caaaccctat ccttctgcca tccagcaaga gccttgccaa caactccacc tttgctcacc 3	53
tccatccct gggttgtcac tctgtgaage ctcgggtccc tgtacttcct gtccgtcccc 4	13
tocageteat tetetteeaa egreggeagee gggaageact tetggetage ettaceeaat 4	
actactcccc actgctttaa atgagaccag ggtccttgtt ttggtgcctt tggatcctat 5	
gatgccttcc cagtctccag ccttggcccc cttctcttct	
atctttcaag tatgtgctac ccaattcctc ttcctcggag gctgctggga cccggaatat 6	
tateccetge tgeaggeete tecaageace acteacetee caggetttee atecgteeca 7	
gtcccaagcc ccatgottca gaacttccct tggcccccc ctacactcca caaattctgg 7	
ggaagtetea enaactgggt ecceteagge ecceaeggga aggaaggtee eccneeaaca 8	333

acntectect gttttecceg gtetecence neegggantt gggeneeena atececaakt 893 tetgaanang aacngeceat tentecentt aaaattaace ttteececee teectgangt 953 958 taggn <210> 38 <211> 81 <212> PRT <213> rat <400> 38 Leu Val Leu Leu Ala Val Ala Leu Gln Thr Ser Asp Ala Gly Pro Tyr -5 -10 Gly Ala Asn Val Glu Asp Ser Ile Cys Cys Gln Asp Tyr Ile Arg His Trp Thr Ser Lys Ser Pro Leu Pro Pro Arg Phe Val Lys Glu Phe Tyr 20 Cys Arg Lys Pro Gly Val Val Leu Ile Thr/Ile Lys Asn Arg Asp Ile Cys Ala Asp Pro Xaa Met Leu Trp Val Lys Lys Ile Leu His Lys Leu 60 Ala <210> 39 <211> 506 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence: S. cerevisiae alpha factor prepro/human MDC cDNA/chimeri/c construct <221> CDS <222> (15) .. (476) <220> <221> mat_peptide <222> (270)..(476) <400> 39 atctcgagct cacg atg /aga ttt cct tca att ttt act gca gtt tta ttc Met/Arg Phe Pro Ser Ile Phe Thr Ala Val Leu Phe -75 -85 -80 gca gca tcc tcc gca tta gct gct cca gtc aac act aca aca gaa gat Ala Ala Ser Ser Ála Leu Ala Ala Pro Val Asn Thr Thr Glu Asp

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-60

-70

			_													1,0030,20
									-22-							<i>/</i> *
													tta Leu			146
													agc Ser			194
													gct Ala			242
													gcc Ala 5			290
													ctg Leu			338
													ccg Pro			386
													gcc Ala			434
									aat Asn 65							476
tga	aggc	ctt	ctag	agcg	gc c	gcat	cgat	a								506
<21 <21 <21	0> 4 1> 1 2> P 3> C 0> 4	54 RT DNA		/			/		/	/	/					
	Arg		Pro	ser	Ile		Thr	Ala	Val	Leu -75		Ala	Ala	Ser	Ser -70	
Ala	Leu	Ala	Ala	P1Q -65		Asn	Thr	Thr	Thr -60		Asp	Glu	Thr	Ala -55		
Ile	Pro	Ala	-50		/val	Ile	Gly	Tyr -45		Asp	Leu	Glu	Gly -40	Asp	Phe	
Asp	Val	Ala -35		. Leu	Pro	Phe	Ser		Ser	Thr	Asn	-25	Gly	Leu	. Leu	
Phe	: Ile -20		Thr	Thr	Ile	-15		lle	· Ala	Ala	Lys -10		Glu	Gly	Val	
					St	JBSTI	TUTE	SHEE	ET (RI	JLE 2	6)					

Pro Leu Asp Lys Arg Gly Pro Tyr Gly Ala Asn Met Glu Asp Ser Val

Cys Cys Arg Asp Tyr Val Arg Tyr Arg Leu Pro Leu Arg Val Val Lys

15 20 25

His Phe Tyr Trp Thr Ser Asp Ser Cys Pro Arg Pro Gly Val Val Leu
30 35 _ 40

Leu Thr Phe Arg Asp Lys Glu Ile Cys Ala Asp Pro Arg Val Pro Trp
45 50 55

Val Lys Met Ile Leu Asn Lys Leu Ser Gln 60 65

<210> 41

<211> 93

<212> PRT

<213> Artificial Human MDC analog

<220>

<223> The amino acid at position 2 is not proline

<220>

<400> 41

Met Ala Arg Leu Gln Thr Ala Leu Leu Val Val Leu Val Leu Leu Ala
-20 -15 -10

Val Ala Leu Gln Ala Thr Glu Ala Gly Xaa Tyr Gly Ala Asn Met Glu
-5 5

Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr Arg Leu Pro Leu Arg

Val Val Lys His Phe Tyr Typ Thr Ser Asp Ser Cys Pro Arg Pro Gly
25 30 35 40

Val Val Leu Leu/Thr Phe Arg Asp Lys Glu Ile Cys Ala Asp Pro Arg
50
55

Val Pro Trp Val Lys Met Ile Leu Asn Lys Leu Ser Gln
60 65

<210> 42

<211> 538

<212> DNA

<213> Homo sapiéms

<220>

<221> CDS

<222> (53)..(\(\beta\)34)

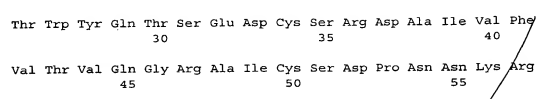
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15



Val Lys Asn Ala Val Lys Tyr Leu Gln Ser Leu Glu Arg Ser

60 65 _ 70

<210> 44

<211> 33

<212> DNA

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence: primer

<400> 44

atgggaccat atggagcaaa tatggaagat agt

33

<210> 45

<211> 335

<212> DNA

<213> Macaque MDC

<220>

<221> CDS

<222> (19)..(297)

<400> 45

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Met Ala Arg Leu Gln Thr Val Phe Leu Gly Val

-20 -15

ctc atc ctc ctt gct gtg gcg ctt caa gca act gag gca ggc ccc tat 99
Leu Ile Leu Leu Ala Val Ala Leu Gln Ala Thr Glu Ala Gly Pro Tyr
-10 5 -1 1

ggc gcc aac atg gaa gac agc gtc tgc tgc cgt gat tac gtc cgt tac 147 Gly Ala Asn Met Glu Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr 5 / 10 15

cgt atg ccc ctg cgt gtg gtg aaa cac ttc tac tgg acc tca gac tcc 195
Arg Met Pro Leu Arg Val Val Lys His Phe Tyr Trp Thr Ser Asp Ser
20 35

tgc ccg agg cct ggc gtg gtg ttg cta acc tcc agg gat aag gag atc 243
Cys Pro Arg Pro Gly Val Val Leu Leu Thr Ser Arg Asp Lys Glu Ile
45 50

tgt gcc gat ccc aga gtg ccc tgg gtg aag atg att ctc aat aag ctg 291 Cys Ala Asp Pro Arg Val Pro Trp Val Lys Met Ile Leu Asn Lys Leu 55 60 65



age caa tgaagageet actatgatga eegtggeeta ageaagee Ser Gln 335

<210> 46

<211> 93

<212> PRT

<213> Macaque MDC .

<400> 46

Met Ala Arg Leu Gln Thr Val Phe Leu Gly Val Leu/Ile Leu Leu Ala
-24
-20
-15
-10

Val Ala Leu Gln Ala Thr Glu Ala Gly Pro Tyr Gly Ala Asn Met Glu

Asp Ser Val Cys Cys Arg Asp Tyr Val Arg Tyr Arg Met Pro Leu Arg
10 15 20

Val Val Lys His Phe Tyr Trp Thr Ser Asp Ser Cys Pro Arg Pro Gly
25 30 40

Val Val Leu Leu Thr Ser Arg Asp Lys Gl/u Ile Cys Ala Asp Pro Arg
45 50 55

Val Pro Trp Val Lys Met Ile Leu Asn/Lys Leu Ser Gln
60 65